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APPLICATION NO.	FILING DATE	FIRST NAMED INVENTOR	ATTORNEY DOCKET NO.	ATTORNEY DOCKET NO. CONFIRMATION NO.	
09/972,203	10/09/2001	Timothy E. Bishop	P 283694 D1056-CIP 3568		
43569 7	590 10/05/2004		EXAMINER		
MAYER, BROWN, ROWE & MAW LLP 1909 K STREET, N.W.			MCCLENDON, SANZA L		
	N, DC 20006		ART UNIT	PAPER NUMBER	
			1711	,	
			DATE MAILED: 10/05/2004		

Please find below and/or attached an Office communication concerning this application or proceeding.

*	Applicati	on No.	Applicant(s)				
	09/972,2	03	BISHOP ET AL.				
Office Action Summary	Examine		Art Unit				
		/IcClendon	1711				
The MAILING DATE of this commun. Period for Reply	ication appears on th	e cover sheet with the o	correspondence addr	ess			
A SHORTENED STATUTORY PERIOD FOR THE MAILING DATE OF THIS COMMUNI  - Extensions of time may be available under the provisions after SIX (6) MONTHS from the mailing date of this comm  - If the period for reply specified above is less than thirty (3)  - If NO period for reply is specified above, the maximum states to reply within the set or extended period for reply Any reply received by the Office later than three months a earned patent term adjustment. See 37 CFR 1.704(b).	CATION. of 37 CFR 1.136(a). In no evunication. D) days, a reply within the state attention period will apply and wwill, by statute. cause the app	ent, however, may a reply be tir utory minimum of thirty (30) day ill expire SIX (6) MONTHS from lication to become ABANDONF	nely filed s will be considered timely. the mailing date of this comi	munication,			
Status							
1) Responsive to communication(s) file	d on 12 July 2004.						
	2b)☐ This action is r	on-final.					
3) Since this application is in condition	• —						
closed in accordance with the practic							
Disposition of Claims							
4) $\boxtimes$ Claim(s) <u>1-29</u> is/are pending in the a	polication.						
4a) Of the above claim(s) is/ai	• •	nsideration					
5)⊠ Claim(s) <u>21 and 22</u> is/are allowed.	•						
6)⊠ Claim(s) <u>1-4,7-20,23 and 26-29</u> is/ar	e rejected.						
7) Claim(s) <u>5,6,24 and 25</u> is/are objected	<del>-</del>						
8) Claim(s) are subject to restric	tion and/or election r	equirement.					
Application Papers							
9) The specification is objected to by the	- Evaminer						
10)⊠ The drawing(s) filed on <u>08 October 2</u>		ented or h) objected	to by the Evaminer				
Applicant may not request that any object							
Replacement drawing sheet(s) including		*	` '	1 121/d)			
11) The oath or declaration is objected to							
Priority under 35 U.S.C. § 119	-			· <del></del> ·			
<u></u>							
12) Acknowledgment is made of a claim f	or foreign priority und	der 35 U.S.C. § 119(a)	-(d) or (f).				
a) ☐ All b) ☐ Some * c) ☐ None of:							
1. Certified copies of the priority							
2. Certified copies of the priority of							
3. Copies of the certified copies of			ed in this National Sta	age			
application from the Internation * See the attached detailed Office action	•	` ''	.d				
ood the ditached detailed Office action	Tion a list of the cent	red copies not receive	a.				
Attachment(s)							
<ol> <li>Notice of References Cited (PTO-892)</li> <li>Notice of Draftsperson's Patent Drawing Review (PTO-892)</li> </ol>	「O-948)	4) Interview Summary Paper No(s)/Mail Da					
3) Information Disclosure Statement(s) (PTO-1449 or F	PTO/SB/08)	5) Notice of Informal Page 1	atent Application (PTO-15	52)			
Paper No(s)/Mail Date <u>9/04</u> .  D.S. Patent and Trademark Office		6)  Other:					
7.5. Patent and Trademark Office PTOL-326 (Rev. 1-04)	Office Action Summa	y Pai	t of Paper No./Mail Date	10012004			

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### DETAILED ACTION

# Response to Amendment

1. In response to the Amendment received on July 12, 2004, the examiner has carefully considered the amendments.

# Response to Arguments

2. Applicant's arguments filed July 12, 2004 have been fully considered but they are not persuasive. Applicant appears to be relying on comparable data in the disclosure to overcome the rejection, drawing attention to example 3, which employs at least 3 photoinitiators (Lucirin<sup>™</sup> TPO, Irgacure<sup>™</sup> 184 and Easacure<sup>™</sup> KIP 100F). However, example 3 is not comparable to comparable exp A (page 43). comparable composition has a different weight amount of the photoinitiator and different compositional components in different weight percent compared to example 3. Applicant's claim 16, states "...said radiation curable composition cures faster than a comparable composition, said comparable composition being identical to said radiation curable composition except that said at least three photoinitiators in said radiation curable composition has been replaced in said comparable composition with an equal weight amount of 1-hydroxy-cyclohexyl-phenyl ketone photoinitiator." Therefore the examiner deems that the showing of unexpected results is not adequately convincing, although is can be seen from the two examples that example 3 has a faster cure, however the data does not positively prove that it is the combination of photoinitiators is the cause for the faster cure. Applicant would need to provide a comparison between two identical compositions except one will have at least three photoinitiator and the other will have an equal weight amount of 1hydroxy·cyclohexyl·phenyl ketone photoinitiator. Since applicant's unexpected result were not persuasive and there are no other convincing arguments that it would have been obvious for an person of ordinary skill level to use a combination of photoinitiators as described in the rejection, the rejections applied in the last Office action still stand.

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# Claim Rejections - 35 USC § 103

3. The following is a quotation of 35 U.S.C. 103(a) which forms the basis for all obviousness rejections set forth in this Office action:

- (a) A patent may not be obtained though the invention is not identically disclosed or described as set forth in section 102 of this title, if the differences between the subject matter sought to be patented and the prior art are such that the subject matter as a whole would have been obvious at the time the invention was made to a person having ordinary skill in the art to which said subject matter pertains. Patentability shall not be negatived by the manner in which the invention was made.
- 4. Claims 1·4, 7·9, 12·20, 23, and 26·29 are rejected under 35 U.S.C. 103(a) as being unpatentable over Snowwhite et al (6,136,880).

Snowwhite et al teaches radiation curable liquid resin compositions for optical fibers. Said liquid resin composition comprises from 10 wt% to 90 wt% of a polyurethane acrylate, from 10wt% to 90wt% of at least one radiation curable monomer diluent and an effective amount of a phosphine oxide photoinitiator having the formula found in the abstract. Said polyurethane acrylate has a molecular weight between 1,000 to about 8,000 Daltons and is the reaction product of an oligomer polyol, a diisocyanate, and a hydroxyl functional ethylenically unsaturated monomer, such as a hydroxyalkyl (meth) acrylate. Said oligomeric polvol is preferably a polyether polyol, such as polytetramethylene glycol—see column 7, lines 10-20 and 50-65. Snowwhite et al teaches, in addition to the preferred polyether polyol, other oligomeric polyols such as polyester, polycarbonate and polyolefin polyols can be used in combination with said polyether polyol. The preferred disocyanates can be chosen from those listed in column 10, lines 27-31, wherein isophorane, methylene bis (4-cyclohexylisocyanate) and toluene diisocyanate are taught. The reactive diluents can be found in column 12, lines 36 to the end. wherein nonylphenol PO modified acrylate is taught. In addition to the above listed reactive diluents Snowwhite et al teaches that di-functional reactive diluents, such as hexanediol diacrylate can be used. Snowwhite et al teaches in addition to the

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phosphine oxide photoinitiator of formula (1) other free radical photoinitiators such as those listed in column 16, lines 35 to 55 can be used in combination with said phosphine oxide. In addition other commonly used additive such as silane coupling agents and others listed in column 17, lines 40.45 can be used in said coating composition. Per examples 1, 5, 7, 8, 9, and 14, Snowwhite et al teaches using polytetramethylene glycol in the synthesis of the polyurethane acrylate. Snowwhite et al teaches using combination of photoinitiators. Therefore it would have been obvious to one skilled in the art at the time of the invention to employ combinations of any photoinitiators taught by Snowwhite et al for the following reasons. Snowwhite et al teaches that in addition to the phosphine oxide other free radical photoinitiators can be used in the radiation curable composition. It is well known in the photopolymerization art to use different kinds of photoinitiators to take advantage of different known properties, such as sensitivity to different wavelengths of light. Acylphosphine oxides, for example, are well known for the ability to provide depth of cure or cure in the presence of pigments and for being useful in combination with phenyl ketone photoinitiators. The motivation for using combination of photoinitiators would have been a reasonable expectation of a faster and more complete cure of said resin compositions in the absence of evidence to the contrary and/or unexpected results. With regard to claim 16, the combination of initiators as taught by Snowwhite et al should inherently cure faster than a comparable composition employing only hydroxy-cyclohexyl-phenyl ketone as a photoinitiator. It is deemed that Snowwhite et al reads on claims 13.16 because Snowwhite et al teaches a composition that reads on the instant composition therefore the compositions of Snowwhite should have the same percentage reacted acrylate unsaturation when cured at a does of about 4.4 mJ/cm2 in the absence of evidence to the contrary.

Note: This application presents comparable data (specifically examples 8-9) to show that the cure rate is improved when a specific combination of photoinitiators employed is used. However claim 16 is not commensurate in scope with the evidence of unexpected results (i.e. example 8 has 5 photoinitiators in an amount of 4.5-wt% and example 9 has four photoinitiators in an amount of 3.0-wt% while the comparative example uses hydroxy-cyclohexylphenyl ketone in amount of 4.0-wt%).

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5. Claims 1·4, 8·9, 12·16, and 19·20 are rejected under 35 U.S.C. 103(a) as being unpatentable over Snowwhite et al (6,359,025).

Snowwhite et al teaches radiation curable liquid resin compositions for optical fiber coatings. Snowwhite et al teaches using a radiation curable urethane acrylate oligomer, a radiation curable monomer diluent and an effective amount of photoinitiator represent by formula 1 found in the abstract. Said polyurethane acrylate has a molecular weight between 1,000 to about 8,000 Daltons and is the reaction product of an oligomer polyol, a diisocyanate, and a hydroxyl functional ethylenically unsaturated monomer, such as a hydroxyalkyl (meth) acrylate. Said oligomeric polyol is preferably a polyether polyol, such as polytetramethylene glycol—see column 7, lines 10.20 and 50.65. Snowwhite et al teaches, in addition to the preferred polyether polyol, other oligomeric polyols such as polyester, polycarbonate and polyolefin polyols can be used in combination with said polyether polyol. The preferred disocyanates can be chosen from those listed in column 10. lines 27-31, wherein isophorane, methylene bis (4-cyclohexylisocyanate) and toluene diisocyanate are taught. The reactive diluents can be found in column 12, lines 36 to the end, wherein nonylphenol PO modified acrylate is taught. In addition to the above listed reactive diluents Snowwhite et al teaches that di-functional reactive diluents, such as hexanediol diacrylate can be used. Snowwhite et al teaches in addition to the phosphine oxide photoinitiator of formula (1) other free radical photoinitiators such as those listed in column 16, lines 35 to 55 can be used in combination with said phosphine oxide. In addition other commonly used additive such as silane coupling agents and others listed in column 17, lines 40.45 can be used in said coating composition. Per examples 1, 5, 7, 8, 9, and 14, Snowwhite et al. teaches using polytetramethylene glycol in the synthesis of the polyurethane acrylate. Snowwhite et al teaches using combination of photoinitiators. Therefore it would have been obvious to one skilled in the art at the time of the invention to employ combinations of any photoinitiators taught by Snowwhite et al for the following reasons. Snowwhite et al teaches that in addition to the phosphine oxide other free radical photoinitiators can be used in the radiation curable composition. It is well known in the photopolymerization art to use different kinds of photoinitiators to take advantage of different known properties, such as sensitivity

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to different wavelengths of light. Acylphosphine oxides, for example, are well known for the ability to provide depth of cure or cure in the presence of pigments and for being useful in combination with phenyl ketone photoinitiators. The motivation for using combination of photoinitiators would have been a reasonable expectation of a faster and more complete cure of said resin compositions in the absence of evidence to the contrary and/or unexpected results. With regard to claim 16, the combination of initiators as taught by Snowwhite et al should inherently cure faster than a comparable composition employing only hydroxy-cyclohexyl-phenyl ketone as a photoinitiator. It is deemed that Snowwhite et al reads on claims 13·16 because Snowwhite et al teaches a composition that reads on the instant composition therefore the compositions of Snowwhite should have the same percentage reacted acrylate unsaturation when cured at a does of about 4.4 mJ/cm2 in the absence of evidence to the contrary.

6. Claims 1, 3-4, 8-9, 13-17 and 18-20 are rejected under 35 U.S.C. 103(a) as being unpatentable over Yamazaki et al (6,057,034).

Yamazaki et al teaches compositions for coating optical fibers comprising a combination of photoinitiators. Per the examples Yamazaki et al teaches reacting polytetramethylene glycol with diisocyanates such as hexamethylene diisocyanate, isophorane diisocyanate and toluene diisocyanate and with a hydroxy alkyl acrylate to obtain a polyurethane acrylate, which anticipates claims 1, 8, and 18-19 and having a molecular weight that read on those found in the claims. The photoinitiators disclosed by applicant, including phenyl ketones, acylphosphine oxide, and oligomeric photoinitiator, re disclosed by Yamazaki et al. See columns 9-10 and the examples, wherein Yamazaki et al specifically mentions commercial combination of photoinitiators in column 10, lines 9-12. Thus it would have been obvious to a skilled artisan at the time of the invention to use combination of any of the photoinitiators taught by Yamazaki et al for the following reasons. Yamazaki et al teaches that of photoinitiators disclosed is useful in the disclosed compositions, the photoinitiators can be used in combination and use of combination of photoinitiators in the examples. It is well known in the photopolymerization art to use different kinds of photoinitiators to take advantage of different known

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properties, such as sensitivity to different wavelengths of light. Acylphosphine oxides, for example, are well known for the ability to provide depth of cure or cure in the presence of pigments and for being useful in combination with phenyl ketone photoinitiators. The motivation for using combination of photoinitiators would have been a reasonable expectation of a faster and more complete cure of said resin compositions in the absence of evidence to the contrary and/or unexpected results. It is deemed that Yamazaki et al reads on claims 13·16 because Yamazaki et al teaches a composition that reads on the instant composition therefore the compositions of Yamazaki et al should have the same percentage reacted acrylate unsaturation when cured at a does of about 4.4 mJ/cm2 in the absence of evidence to the contrary.

7. Claims 1, 3, 8, and 10·20 are rejected under 35 U.S.C. 103(a) as being unpatentable over Moschovis et al (4,782,129).

Moschovis et al teaches buffer coating compositions for optical fibers. Said composition comprises an acrylated capped polyurethane in admixture with a monoethylenically unsaturated monomer. Said polyurethane is the reaction product of an organic diisocyanate with a modified diol and a hydroxyalkyl acrylate. Said modified diol is the diester reaction product of polyoxytetrmethylene glycol with a long chain ester forming hydrocarbon substituted dicarboxylic acid or a ester forming substitution product thereof. The dicarboxylic acid can be succinic acid or adipic acid. The organic diisocyanate can be a toluene diisocyanate, isophorane diisocyanate, a dimer fatty acid diisocyanate and, as seen in the examples, saturated methylene diphenyl diisocyanate, wherein some are deemed to read on those found in the claims. In addition to the modified diol, other diols, such as 1,6 hexane diol and trimethylolpropane in amounts up to 10% of the total amount of difunctional reactants in the polyurethane, can be used for making the polyurethane. It is deemed that applicant's component proportions are read by the general disclosure as taught by Moschovis in the absence of arguments to the contrary. Per examples, Moschovis et al teaches using a silane coupling agent (e.g., methacrylate ester of gamma hydroxylpropyl trimethoxy silane). Moschovis et al teaches using photoinitiators as a component in said compositions. Moschovis et al teaches using photoinitiator combination of Irgacure 907 with isopropylthioxanthone, in addition also phosphine oxides and hydroxylcyclohexyl phenyl ketone are also useful.

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Moschovis et al teaches that said photoinitiators could be used alone or in admixture with one another. Thus it would have been obvious to a skilled artisan at the time of the invention to use combination of any of the photoinitiators taught by Moschovis et al for the following reasons. Moschovis et al teaches that of photoinitiators disclosed is useful in the disclosed compositions, the photoinitiators can be used in combination and use of combination of photoinitiators in the examples. It is well known in the photopolymerization art to use different kinds of photoinitiators to take advantage of different known properties, such as sensitivity to different wavelengths of light. Acylphosphine oxides, for example, are well known for the ability to provide depth of cure or cure in the presence of pigments and for being useful in combination with phenyl ketone photoinitiators. The motivation for using combination of photoinitiators would have been a reasonable expectation of a faster and more complete cure of said resin compositions in the absence of evidence to the contrary and/or unexpected results. It is deemed that Moschovis et al reads on claims 13-16 because Moschovis et al teaches a composition that reads on the instant composition therefore the compositions of Moschovis et al should have the same percentage reacted acrylate unsaturation when cured at a does of about 4.4 mJ/cm2 in the absence of evidence to the contrary.

# Allowable Subject Matter

- 8. Claims 5·6 and 24·25 are objected to as being dependent upon a rejected base claim, but would be allowable if rewritten in independent form including all of the limitations of the base claim and any intervening claims.
- 9. The following is a statement of reasons for the indication of allowable subject matter: The prior art fails to teach a radiation curable oligomer having the individual proportion amounts found in claims 5.6 and 25. Nor using a polydimethyl siloxane in a composition as taught in claim 23.
- 10. Claims 21.22 are allowed.
- 11. The following is an examiner's statement of reasons for allowance: The primary reasons for allowance is the radiation curable oligomer comprising from 10-30 wt% of isophorane diisocyanate, 5·15 wt% of dicyclohexylemthane diisocyanate,

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45.75 wt% of polytetramethylene glycol and 5.20 wt% of hydroxyethylacrylate, which when cured at a dose of about 4.4 mJ/cm2 has a percentage reacted acrylate unsaturation of at least 60%.

Any comments considered necessary by applicant must be submitted no later than the payment of the issue fee and, to avoid processing delays, should preferably accompany the issue fee. Such submissions should be clearly labeled "Comments on Statement of Reasons for Allowance."

### Conclusion

12. THIS ACTION IS MADE FINAL. Applicant is reminded of the extension of time policy as set forth in 37 CFR 1.136(a).

A shortened statutory period for reply to this final action is set to expire THREE MONTHS from the mailing date of this action. In the event a first reply is filed within TWO MONTHS of the mailing date of this final action and the advisory action is not mailed until after the end of the THREE MONTH shortened statutory period, then the shortened statutory period will expire on the date the advisory action is mailed, and any extension fee pursuant to 37 CFR 1.136(a) will be calculated from the mailing date of the advisory action. In no event, however, will the statutory period for reply expire later than SIX MONTHS from the mailing date of this final action.

13. Any inquiry concerning this communication or earlier communications from the examiner should be directed to Sanza L McClendon whose telephone number is (571) 272·1074. The examiner can normally be reached on Monday through Friday 7:30-4:00.

If attempts to reach the examiner by telephone are unsuccessful, the examiner's supervisor, James Seidleck can be reached on (571) 272-1078. The fax phone number for the organization where this application or proceeding is assigned is 703-872-9306.

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Sanza L McClendon Examiner

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SMc

James J. Seidleck
Supervisory Patent Examirant
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